
Ultrafast dynamics in van der Waals heterostructures

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Abstract

Van der Waals heterostructures are synthetic quantum materials composed of stacks of atomically thin two-dimensional (2D) layers. Because the electrons in the atomically thin 2D layers are exposed to layer-layer coupling, the properties of van der Waals heterostructures are defined not only by the constituent monolayers, but also by the interactions between the layers. Many fascinating electrical, optical, and magnetic properties have recently been reported in different types of van der Waals heterostructures. In this review we focus on unique excited-state dynamics in transition metal dichalcogenide (TMDC) heterostructures. TMDC monolayers are the most widely studied 2D semiconductors, featuring prominent exciton states and accessibility to the valley degree of freedom. Many TMDC heterostructures are characterized by a staggered band alignment. This band alignment has profound effects on the evolution of the excited states in heterostructures, including ultrafast charge transfer between the layers, the formation of interlayer excitons, and the existence of long-lived spin and valley polarization in resident carriers. Here we review recent experimental and theoretical efforts to elucidate electron dynamics in TMDC heterostructures, extending from time scales of femtoseconds to microseconds, and comment on the relevance of these effects for potential applications in optoelectronic and valleytronic/spintronic devices.

36 **Main text**

37 Advances in the isolation and manipulation of atomically-thin sheets of two-dimensional (2D) crystals,
38 starting with the investigations of graphene a decade ago, have ushered in a new era of basic
39 scientific research and technological innovation. 2D layers with diverse properties can now be
40 prepared separately and then stacked together to form new types of quantum materials, known as
41 van der Waals (vdW) heterostructures. The ability to combine materials with monolayer precision
42 enables the design and creation of functional 2D materials that do not exist in nature. Today we have
43 at our disposal a wide variety of atomically thin 2D layers, ranging from semiconducting MoS₂ and
44 insulating hexagonal boron-nitride (h-BN) to magnetic CrI₃ and superconducting NbSe₂, that can be
45 stacked one upon the other. Since the electrons in atomically thin layers are exposed, different
46 quantum states found in the individual layers can interact and couple to one another in ways that are
47 not possible in other systems.

48
49 VdW heterostructures constitute a vast family of new quantum materials, since they are defined not
50 only by the combination of constituent monolayer materials, but also by the stacking sequence and
51 relative crystallographic alignment of the layers. Further control of physical properties in 2D vdW
52 heterostructures can be achieved through the application of electrostatic gating and fields, as well as
53 substrate and strain engineering. Many fascinating physical phenomena have been reported in
54 different vdW heterostructures, as exemplified by transport measurements revealing Hofstadter
55 butterfly states, fractional Chern insulators, gate-tunable Mott insulators, unconventional
56 superconductivity, etc¹⁻⁷. In addition to electrical transport, there has also been great progress in the
57 study of the optical properties and excited-state dynamics in vdW heterostructures. Here we review
58 the new dynamical phenomena that emerge in semiconducting vdW heterostructures composed of
59 stacked transition metal dichalcogenide (TMDC) layers. We focus our discussion on TMDC
60 heterostructures, since the individual TMDC layers, with their many distinctive and intriguing
61 properties, have already been well characterized and provide a strong basis for understanding the
62 emergent new properties of heterostructures.

63
64 TMDC semiconductors (MX₂ layers with 2H symmetry and M = Mo, W; X = S, Se, Te) exhibit direct
65 gaps at monolayer thickness. They feature strong light-matter interactions and dramatically enhanced
66 electron-electron interactions, with the optical properties largely defined by exciton states. The
67 exciton binding energies in monolayer TMDCs are hundreds of meV— as much as two orders of
68 magnitude larger than in typical bulk semiconductors like silicon or GaAs^{8,9}. In addition, TMDC
69 monolayers provide a platform to investigate and control the valley degree of freedom — often
70 designated as valley pseudospin — associated with the energetically degenerate K and K' valleys and
71 accessible optically through the presence of valley circular dichroism. The valley pseudospin in TMDC
72 is, moreover, coupled to the electron/hole spin due to strong spin-orbit interactions^{10,11}.

73
74 Understanding the dynamic interplay and evolution of the charge, spin, and valley excitations in vdW
75 heterostructures is of fundamental scientific interest. It is also of central importance for many potential
76 applications of TMDC materials in optoelectronics, spintronics, and valleytronics. The dissociation of
77 optically excited excitons into free carriers is, for example, critical for photovoltaic devices, while the
78 ability to control and stabilize valley polarization is essential for valleytronic applications. The
79 formation of vdW heterostructures in TMDC layers can profoundly affect their excited-state dynamics,
80 ranging from the dissociation of intralayer excitons and the formation of interlayer excitons to the
81 relaxation of spin and valley polarization. The use of vdW heterostructures provides a powerful
82 platform to control and optimize the dynamic response of the constituent materials. In this review, we

83 survey recent progress in probing electron dynamics, extending from femtoseconds to microseconds,
84 in TMDC heterostructures. On short time scales ($\lesssim 1$ ps), the dynamics is dominated by the charge
85 transfer and energy relaxation processes in the heterostructure. On longer time scales ($\gtrsim 1$ ps), the
86 recombination of interlayer excitons and relaxation of the spin and valley degrees of freedom become
87 relevant. The rate of these processes can vary by orders of magnitude depending on the configuration,
88 temperature, and doping of the heterostructure. We will also touch upon the dynamics of lateral
89 transport of spin and valley polarization in TMDC heterostructures. We describe the physical
90 mechanisms that underlie the different types of dynamic response in TMDC heterostructures and
91 distinguish the behavior from that of the constituent layers, as well as mention briefly the implications
92 for potential new technologies.

94 **Band alignment in TMDC heterostructures**

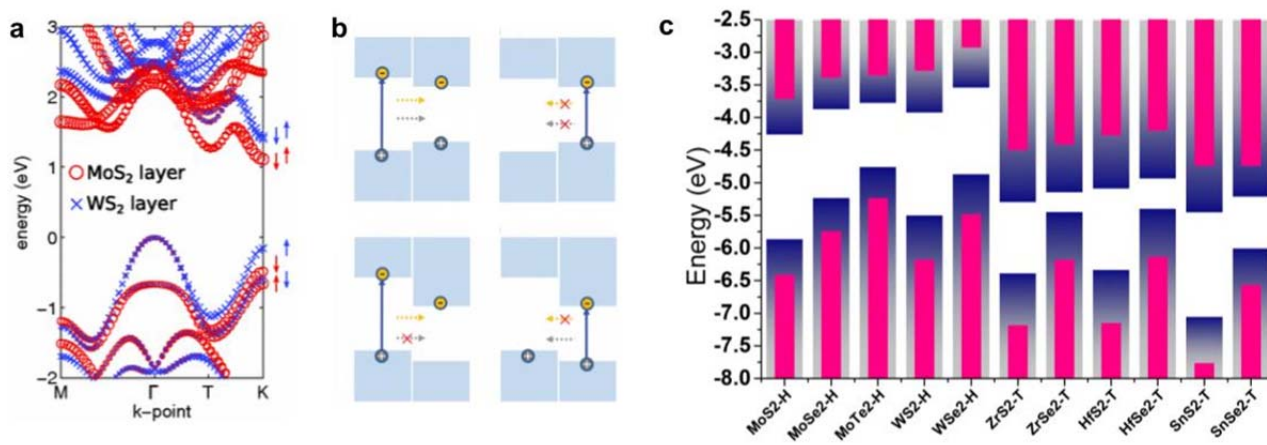
95 In describing the single-particle electronic states in a vertical TMDC heterostructure, most authors in
96 the literature consider the states to be largely localized within the individual layers. This approximation
97 holds well for electronic states close to the band gap due to unusually weak interlayer coupling in
98 TMDCs at the relevant K and K' points in the Brillouin zone (Fig. 1a)¹². Therefore one can directly
99 examine the relative energy difference, or band alignment, of material-specific electronic band
100 extrema, just as is commonly done for heterostructures and quantum wells based on bulk
101 materials^{13,14}. An important distinction for atomically thin layers is that the concept of band bending
102 near an interface does not apply¹⁵.

104 Semiconductor heterostructures can have different types of band alignment depending on the energy
105 difference of the conduction and valence band extrema in the constituent layers (Fig. 1b)^{13,14}. The
106 lowest-lying states for electrons and holes reside in the same layer for a type-I heterostructure¹⁶, but
107 are separated in different layers for a type-II heterostructure^{13,17}. Further offsets in energy could, in
108 principle, lower the conduction band minimum (CBM) of one layer to a position below the valence
109 band maximum (VBM) of the other layer, yielding a type-III heterostructure¹⁸.

111 The type of band alignment has profound effects on the excited-state dynamics in a vdW
112 heterostructure. In a type-I heterostructure carriers can only flow from the layer with the larger band
113 gap to the layer with the smaller band gap (assuming that they do not have excess energy greater
114 than the band offsets). The same applies to energy transfer through the exchange of virtual photons.
115 In a type-II heterostructure, on the other hand, electrons accumulate in the layer with the lower CBM,
116 while holes accumulate in the other layer^{17,19,20}. Energy transfer can, however, still occur from the
117 larger-gap to the smaller-gap material²¹.

119 There have been numerous theoretical studies of the band alignment in TMDC
120 heterostructures^{13,14,18,22-24}. The result of one recent calculation is shown in Fig. 1c. Many of the
121 theoretical studies make use of DFT or related methods and cannot accurately account for the effects
122 of doping (Fermi level difference), dielectric screening, and excitonic interactions²⁵. They provide
123 nonetheless guidelines for realizing specific types of heterostructures and combination of band gaps.
124 Although theoretical studies are advancing rapidly, at present a definitive determination of
125 heterostructure band alignment must rely on experiment^{26,27}.

127 In the following, we describe experimental findings concerning the dynamics of charge and energy
128 transfer for representative TMDC heterostructures, followed by a summary of the relevant theoretical
129 studies.
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131
132 **Fig. 1 Band alignment in vertical vdW heterostructures of TMDCs.** (a)
133 Calculated electronic states in a MoS₂/WS₂ vertical heterostructure,
134 showing layer-localized states near the band edges at the K point¹². (b)
135 Schematic of allowed charge transfer in heterostructures with type-I (top)
136 and type-II (bottom) band alignment¹⁶. (c) Calculated band-edge energies
137 for various TMDCs²² based on different theoretical treatments: DFT-PBE
138 (blue) and G₀W₀ (pink).

139 Charge transfer and energy transfer in TMDC heterostructures - Experiment

140 Following the predictions of type-II band alignment in TMDC heterostructures, several experimental
141 studies were reported that aimed (among other goals) to validate these theoretical results by probing
142 the associated charge transfer (CT) or energy transfer (ET) processes. Vertical heterostructures have
143 been examined for various material combinations prepared with different fabrication methods,
144 including stacking of layers exfoliated from bulk crystals and from layers grown by chemical vapor
145 deposition (CVD), as well as heterostructures grown directly by CVD.

146
147 Ultrafast optical measurements using pump-probe spectroscopy provide the possibility of accessing
148 charge and energy transfer processes with femtosecond (fs) time resolution. In initial experiments,
149 Hong *et al.* examined the dynamics of the MoS₂/WS₂ heterostructure. Following excitation by an
150 ultrafast laser pulse resonant with the lower-energy MoS₂ A exciton, they observed a transient change
151 in reflectivity near the *higher-energy* WS₂ exciton (Fig. 2a)¹⁹. Based on theoretical predictions of type-
152 II band alignment of the two materials, this transient response was identified as arising from CT of a
153 hole from the MoS₂ monolayer to the WS₂ monolayer. By deconvolving the instrument response from
154 the rise-time of the signal, the authors were able to provide an upper limit of 50 fs for the charge
155 transfer time. Ceballos *et al.* observed similar dynamics in a heterostructure of MoS₂/MoSe₂, without
156 spectrally resolving the reflection of the probe²⁸. They concluded, based on the theoretically predicted
157 band alignment, that electrons were transferred from MoSe₂ to MoS₂. In addition, when exciting both
158 layers (using excitation resonant with the higher-energy exciton feature in MoS₂) and comparing the

159 transient reflectivity signal of the monolayers to that of the heterostructure, the authors identified the
160 presence of hole transfer in the opposite direction, *i.e.*, from MoS₂ to MoSe₂ (Fig. 2b).

161

162 Following these first experimental investigations, several groups examined the nature of the ultrafast
163 CT for different types of heterostructures under different conditions. Heo *et al.* compared CVD-grown
164 and manually stacked WS₂/MoS₂ heterostructures with a focus on the dependence on the relative
165 (twist) angle of the two constituent lattices²⁹. Measuring the transient transmission of the probe signal,
166 they did not resolve differences in the rise time of the signal associated with CT, but did observe a
167 pronounced difference in the decay times of the signal. Wang *et al.* examined several WSe₂/WS₂
168 heterostructures composed of CVD-grown layers mechanically stacked with different twist angles^{20,29}.
169 Using pump-probe measurements, together with steady-state techniques such as reflection contrast
170 spectroscopy, they showed the same ultrafast signature from the onset of CT, either from electrons
171 moving from WSe₂ to WS₂ or from holes traveling in the opposite direction. They concluded that
172 interlayer CT takes place within 450 fs, close to the duration of the pump pulses in their experiment,
173 and does not exhibit measurable sensitivity to the twist angle. Further reinforcing this conclusion, Zhu
174 *et al.* explored deterministically aligned heterostructures of mechanically exfoliated MoS₂ and WSe₂;
175 they that the CT signal appears within 40 fs regardless of twist angle (Fig. 2c)³⁰. On the other hand,
176 the time scale of the decay was varied with the twist angle, but without any clear trend. Ji *et al.*
177 reported a similar CT rise time for stacks of CVD-grown MoS₂ and WS₂ (Ref³¹). Such rapid interlayer
178 CT irrespective of crystal orientation (and thus crystal momentum) is somewhat unintuitive. Chen *et*
179 *al.* probed intraband transitions in a heterostructure using infrared light and suggested a potential
180 explanation based on the rapid formation of “hot” interlayer excitons³². Additional theoretical
181 investigations are summarized in the next section.

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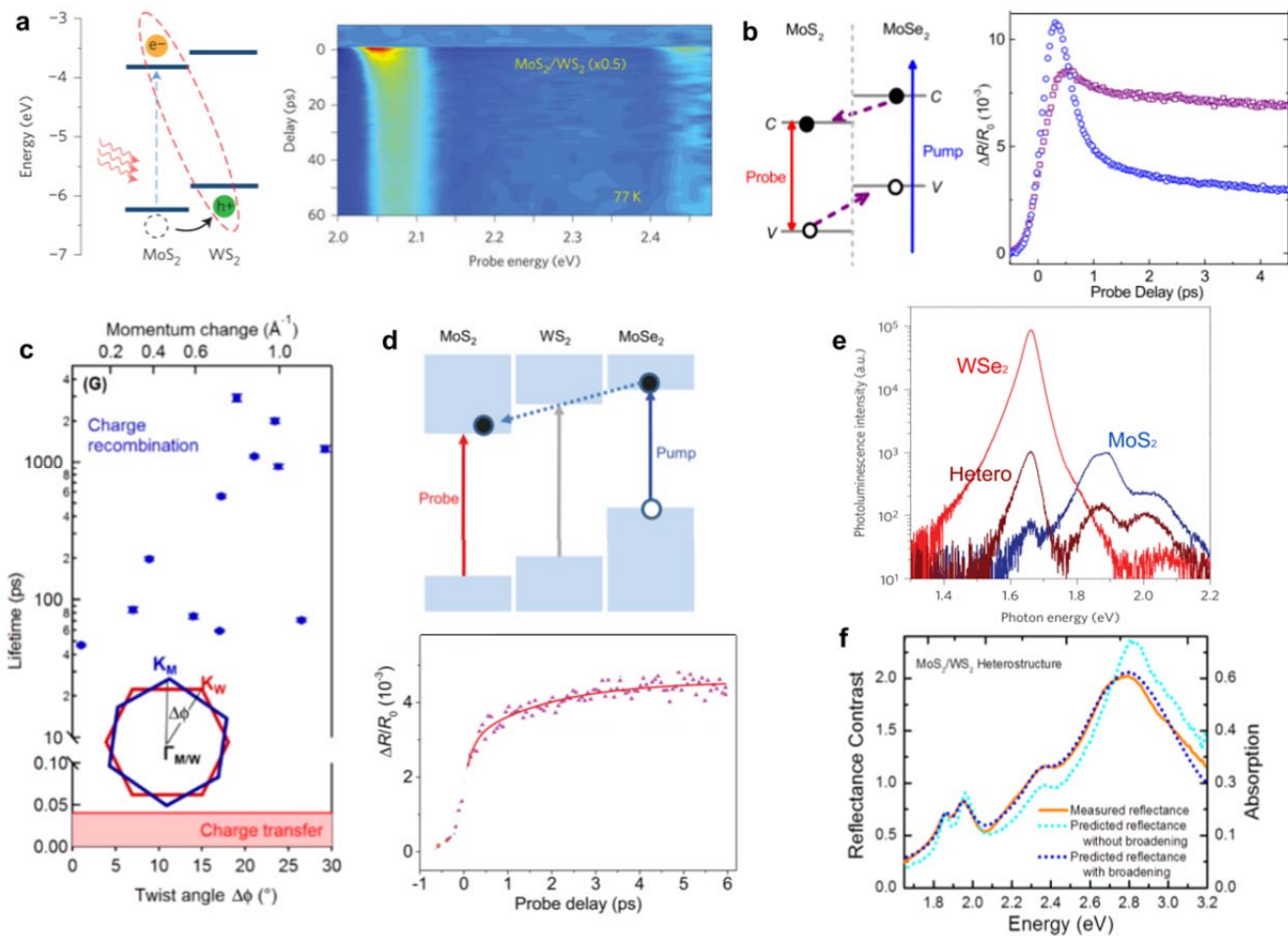


Fig. 2 Experimental studies of ultrafast charge transfer in vertical TMDC heterostructures. (a) Schematic and energy-resolved transient absorption spectra of a MoS₂/WS₂ heterostructure, excited by an optical pulse near the lower-energy MoS₂ A exciton feature, indicating hole transfer¹⁹. (b) Schematic and time-resolved differential reflection of a MoS₂/MoSe₂ heterostructure (blue) and of monolayer MoS₂ (purple), excited by an optical pulse above the band gaps of both materials, indicating both electron and hole transfer²⁸. (c) Charge transfer (red shading) and recombination (blue) times in heterostructures of MoS₂ and WSe₂ with different twist angles, as indicated³⁰. (d) Differential reflection at the energy of the MoS₂ A exciton following excitation at the energy of the MoSe₂ A exciton in a MoSe₂/WS₂/MoS₂ trilayer heterostructure³³. (e) PL quenching³⁴ in MoS₂/WSe₂ and (f) broadening of the features absorption features³⁵ in a MoS₂/WS₂ heterostructure compared to the response of the separated monolayers.

Further demonstrating its robustness, the signature of ~ps electron transfer was observed in a trilayer heterostructure where monolayers of MoSe₂ and MoS₂ were separated by a monolayer WS₂ (Fig. 2d, Ref³³). The authors suggested that such rapid CT across multiple materials is coherent, rather than sequential.

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205 CT dynamics was also revealed in investigations of the coupling between pairs of TMDC layers. CT in
206 a MoS₂/MoTe₂ heterostructure was identified using pump-probe experiments³⁶, leading to the
207 suggestion of MoTe₂ as a good electrical contact for other semiconducting TMDCs and metals. Later,
208 cascaded transfer of electrons and holes across several TMDCs heterostructures was invoked to
209 explain correlated blinking observed in the photoluminescence (PL) in those stacks³⁷.

210
211 In addition to these time-resolved pump-probe measurements, CT between two TMDC layers in a
212 heterostructure has been inferred from quenching of the photoluminescence of the constituent
213 monolayers^{19,20,28-33,38-40} and by broadening of the resonant features in the monolayers in optical
214 absorption measurements^{20,35}, both phenomena arising as a consequence of the presence of
215 additional relaxation channels in the heterostructures. A reduction in the PL intensity by factor of a few
216 tens to a few hundreds has commonly been observed (Fig. 2e)^{19,34}. This suggests a corresponding
217 ratio for the charge transfer time compared to the population lifetime in the isolated material.
218 Assuming the latter to be a few hundred picoseconds for excitons in monolayer TMDCs⁴¹ at room
219 temperature, we estimate a CT time of ~1 ps, somewhat longer than measured by pump-probe
220 techniques. This discrepancy can be explained as the result of a small fraction of the heterostructure
221 exhibiting poor contact between the layers, thus yielding reduced PL quenching compared to that of
222 the ideal structure. In optical absorption measurements, linewidth broadening of heterostructures
223 compared with that of the separate monolayers has been reported and used to estimate non-radiative
224 decay rates comparable to those deduced from pump-probe experiments (Fig. 2f)³⁵. Extrinsic factors,
225 such as strain introduced in fabricating the heterostructure, can potentially also play a role. The
226 extremely rapid (few femtosecond) relaxation times inferred for high-lying states may, however,
227 remain difficult to probe directly in the time domain, but is easily observable by lineshape analysis.

228
229 Another dynamic process that may compete with CT is energy transfer (ET). In this latter process, an
230 exciton created in one layer recombines, and the released energy creates an exciton in the other layer.
231 Kozawa *et al.* reported evidence for such a process in a MoSe₂/WS₂ heterostructure (Ref²¹) on the
232 basis of a measured enhancement of PL from the MoSe₂ feature under excitation resonant with the
233 WS₂ optical band gap. Recently, ET was also identified in heterostructures of MoS₂ and WS₂
234 separated by insulating layers of h-BN⁴². In these structures, PL quenching from the heterostructure
235 was reduced, or even become PL enhancement, upon increasing the thickness of the h-BN spacer.
236 This was interpreted as an ET process between B excitons in MoS₂ and A excitons in WS₂. The
237 dependence of the enhancement on spacer thickness, with an optimum of PL enhancement for ~5
238 layers of h-BN and subsequent reduction of this effect with increasing layer thickness, is compatible
239 with the predicted trend for a dipole-dipole interaction. This highlights the major difference between
240 ET and CT processes: While the latter requires intimate coupling of the two constituent monolayers,
241 the former, originating in dipole-dipole interactions, can act at greater distances and across insulating
242 spacers. Although this difference between the processes is clear, there is currently little direct
243 experimental information on the absolute rates of energy transfer for TMDC layers and in what
244 regimes and under what conditions energy transfer competes with charge transfer.

245 **Charge transfer and energy transfer in TMDC heterostructures – Interpretation and theory**

246 Since most experiments have excited and probed excitonic resonances of the constituent layers, it
247 has often been assumed that CT occurs between their direct band-edge (K/K' valley) states. Band
248 structure calculations have shown that the K/K' states are localized around the central layer of metal
249 atoms and have weak interlayer interactions. On the other hand, for states of different momentum,

250 such as in the Γ or Q valleys, the interlayer coupling may be significantly stronger. One consequence
251 of this difference is the transition from indirect to direct band gap upon thinning TMDCs to the
252 monolayer limit: the bulk CBM in the Q valley, more affected by interlayer coupling, lies above the K
253 valley in monolayers^{43,44}. For the same reason, the K-valley states are not expected to show such
254 rapid interlayer charge transfer. This contradiction with experiment is further heightened by the
255 apparent independence of CT on twist angle and lattice mismatch, the factors that dictate the
256 momentum difference between the initial and final states, as well as its insensitivity to temperature.
257 Here we briefly survey some of the approaches presented in the literature to identify the mechanism
258 responsible for the very efficient CT processes observed experimentally in vertical TMDC
259 heterostructures.

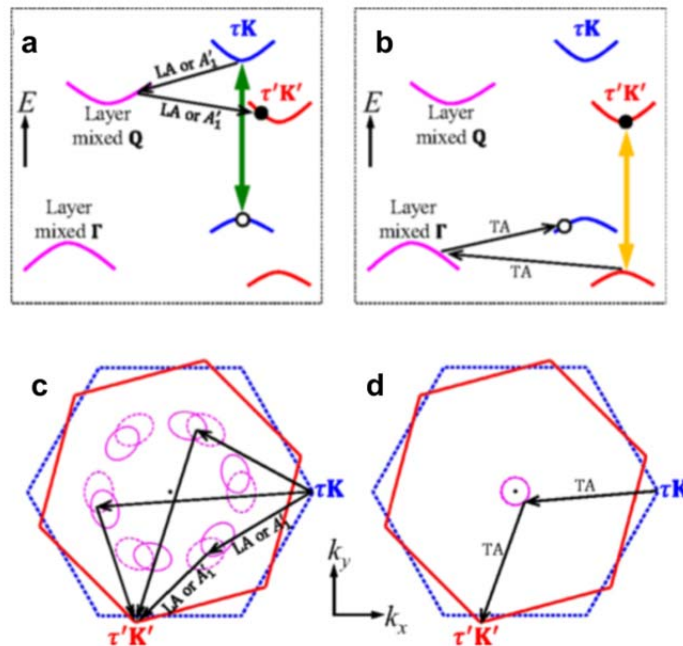
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261 Zhu *et al.*, following considerations relevant for CT processes in molecular systems, have pointed out
262 the possible role of localization in bridging the momentum mismatch⁴⁵. The electrostatic attraction
263 between the optically excited electron and hole, which leads to the formation of tightly bound excitons,
264 gives rise to a distribution of momenta for the charge carriers across a significant range wavevectors
265 in the Brillouin zone (BZ). This situation could explain the apparent lack of momentum conservation in
266 the observed CT process. However, it is unclear with this effect alone could supply the large
267 momentum required for CT in heterostructures with large twist angles³⁰. In addition, for excitation
268 substantially above the band gap, it is unclear whether exciton formation occurs prior to charge
269 transfer²⁸.

270
271 Several groups have also employed numerical calculations using molecular dynamics (MD) and time-
272 dependent density-functional theory (TD-DFT) to elucidate the mechanism for interlayer charge
273 transfer. Since accurately accounting for the effect of excitons is computationally demanding, these
274 calculations have focused on the transfer of free charge carriers from one monolayer to its neighbor.
275 Wang *et al.* described a process of transferring holes directly between the K valleys of the two layers
276 mediated by the electric dipole interaction of the initial and final states, which enhances the coupling
277 between the states to above a critical level for collective charge transfer⁴⁶. In this description, both
278 twist angle and temperature have a significant influence on the transfer rate⁴⁶. Zhang *et al.* pointed
279 out the significance of such dipole coupling between specific states in the vicinity of the K valley and
280 highlighted the expected twist-angle dependence of the CT rate⁴⁷. Additionally, the authors argue that
281 the omission of excitonic effects for above gap excitation may not be significant if the time scales for
282 CT and exciton formation are comparable. (This argument is, however, problematic for explaining CT
283 from excitons created directly by resonant excitation.)

284
285 A slightly different picture is presented by Long *et al.*⁴⁸ and by Li *et al.*⁴⁹ for CT processes in
286 MoSe₂/MoS₂ and MoS₂/WS₂ systems, respectively. They propose that holes and/or electrons undergo
287 transfer from one layer to the other due to mixing between the electronic states near the K point of
288 both layers^{48,49}: the states into which the charge carriers are optically injected are coherently mixed
289 (and therefore delocalized) across the two layers, so that CT need only to be driven by an intralayer
290 relaxation process to the K-valley, mediated by phonon emission. In this model, the coupling relies on
291 specific layer orientations to facilitate the state mixing in the heterostructure. The discrepancy
292 between this requirement and the apparent twist-angle independence of CT in experiments is
293 explained in terms of changes in the relative local atomic positions in the two layers across the moiré
294 pattern of the heterostructure^{50,51}. Regardless of the twist angle of the layers, such lateral
295 inhomogeneity provides regions where the coupling between the layers is strong³¹. In all of the above
296 scenarios, phonons are necessary for the completion of the CT process, but not for its initiation.

297

298 Recent work has also explored a more direct role for phonons in the initiation of CT. Wang *et al.*
299 considered momentum-conserving charge transfer between the layers in various regions of the BZ,
300 including at the K, Q and Γ points for different twist angles and degrees of lattice mismatch⁵², as
301 shown in Fig. 3a-d. The proposed mechanism involves scattering by phonons from the K valley to the
302 Γ valley (for holes) or to the Q valley (for electrons), regions where the interlayer coupling is strong
303 and interlayer charge transfer is rapid. After charge transfer, scattering with another phonon would
304 bring the charge carriers back to the K-valley. For intralayer scattering within 20 fs, the entire charge
305 transfer process would occur in less than 100 fs, as observed experimentally. A similar scheme was
306 developed by Zheng *et al.* using a numerical MD TD-DFT calculation, although without accounting for
307 inhomogeneities from the moiré pattern⁵³. These theories better match the observed twist-angle
308 independence, as the coupling around the Γ and Q points is not sensitive to the layer orientation. The
309 expected temperature dependence of this mechanism is also weak, as it only requires emission of
310 phonons to dissipate the excess energy available from the transfer. This general mechanism has
311 been adopted in the interpretation of recent experiments related to CT processes^{30,33,39}.
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316 **Fig. 3 Theoretical concepts explaining the robust and ultrafast**
317 **nature of CT in TMDC heterostructures⁵².** (a) Schematic representation
318 of phonon mediated electron transfer. (b) The same for hole transfer. (c)
319 Top view of the twisted BZ of the two layers, where electron transfer
320 between the K-points is mediated by phonon coupling to the Q-point. (d)
321 The same for hole transfer, mediated in this case by the coupling to the
322 Γ -point.

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324 **Spin and valley dynamics in TMDC heterostructures**

325 The previous sections addressed carrier dynamics in TMDC heterostructures on the ultrafast time
326 scale (≤ 1 ps) relevant for interlayer charge transfer in systems with type-II band alignment. In the
327 following section, we focus on spin and valley relaxation dynamics in TMDC heterostructures, which
328 take place on considerably longer time scales.

329
330 Within a TMDC monolayer, there are two distinct relaxation processes to consider. First, the
331 population decay of optically excited excitons has a characteristic time scale of few picoseconds to
332 nanoseconds, depending on choice of material, sample preparation, temperature, etc.⁵⁴⁻⁵⁸. Second,
333 the exciton spin-valley lifetime, which determines how long information can be stored in the spin-
334 valley degree of freedom, has been found to be a few picoseconds in isolated monolayer TMDCs⁵⁹⁻⁶³.
335 Both the population and the spin-valley lifetimes in type-II heterostructures, where the electrons and
336 holes reside in different layers after the rapid initial charge transfer process, can differ markedly from
337 the corresponding lifetimes in isolated monolayers.

338
339 Below we review recent experimental results on dynamics on the pico-to-micro-second time scale in
340 heterostructures and show that both the population lifetime and the spin-valley lifetime can be
341 significantly longer than for the monolayer case. We then summarize recent efforts towards
342 understanding the physical mechanisms of the corresponding intervalley scattering processes. Finally,
343 we discuss the spatio-temporal dynamics of spin and valley polarization in TMDC heterostructures.
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345 **Long spin and valley lifetime in TMDC heterostructures**

346 Traditional electronic devices are based on the manipulation of electron charges in the real space.
347 Using other electron degrees of freedom as the information carrier, such as spin and valley, can
348 potentially overcome fundamental limits of speed and power consumption, giving rise to intriguing new
349 concepts in spintronics and valleytronics. A long spin/valley lifetime is necessary to ensure that the
350 spin/valley information will be maintained in the idle state and can persist long enough to be
351 processed. We note that the valley lifetime discussed here should not be confused with the valley
352 depolarization time: the former can originate from different mechanisms, including the population
353 decay of the valley information carriers, while the latter only describes the intervalley scattering
354 process.

355
356 TMDCs offer a promising platform for spintronic and valleytronic applications, owing to several
357 attractive properties of these materials. The valley-dependent optical selection rule allows for
358 convenient creation, manipulation, and detection of excitons in specific valleys with circularly polarized
359 light⁶⁴⁻⁶⁶. Furthermore, the spin-valley locking effect suggests that a very long spin-valley lifetime is
360 possible because intervalley scattering of electrons or holes requires both a large momentum change
361 from K to K' and a simultaneous spin flip^{10,11}. However, several groups have measured exciton spin-
362 valley dynamics using time-resolved Kerr rotation (TRKR)⁵⁹⁻⁶³, and the spin-valley lifetime was found
363 to be rather short, ranging from one to few picoseconds, even at low temperatures. This counter-
364 intuitive observation was later explained as a consequence of the exchange interaction between
365 excitons in the two valleys⁶⁷⁻⁷⁰: a bright exciton always has total momentum and spin of zero and
366 therefore does not require any change in momentum or spin to scatter to the other valley as an intact
367 exciton. This reduced spin-valley lifetime of excitons in TMDC monolayers significantly limits their use
368 in carrying spin-valley information.

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370 A general strategy for improving the spin-valley lifetime in TMDCs is to eliminate the exciton exchange
371 interaction by converting excitons into other excitations that serve as alternative carriers of valley
372 information. An additional figure of merit, the conversion efficiency, must be introduced to quantify the
373 final valley imbalance created from each optically excited exciton. To avoid loss of valley information
374 during conversion, the time scale of the conversion process (*i.e.*, the exciton population lifetime) must
375 be comparable to or shorter than the picosecond spin-valley lifetime of excitons. In monolayer TMDCs,
376 several candidates have been considered as the alternative information carriers, including trions⁷¹⁻⁷³,
377 dark excitons^{74,75}, biexcitons^{76,77}, and resident carriers⁷⁸⁻⁸¹. For example, Fig. 4a,b show the valley
378 dynamics of trions and resident electrons probed by TRKR measurement. Unlike bright excitons,
379 these excitations have non-zero total momentum and/or total spin, and therefore will not suffer from
380 rapid spin-valley relaxation through the exchange interaction. Their spin-valley lifetimes at low
381 temperatures range from tens of picoseconds to a microseconds, but the conversion efficiency from
382 the initially generated exciton has rarely been characterized. However, since the exciton population
383 lifetime is comparable to or longer than the exciton spin-valley lifetime in these cases, the valley
384 conversion efficiency is likely to be considerably less than unity.

385

386 On the other hand, the interlayer charge transfer process in type-II heterostructures provides an
387 attractive mechanism to break intralayer excitons on the femtosecond time scale. As discussed above,
388 the ultrafast charge transfer process occurs very rapidly, typically within ~50 fs, in TMDC
389 heterostructures^{19,82}. Because this time scale is much faster than exciton spin-valley relaxation, the
390 loss of spin-valley information during the conversion process is expected to be minor.

391

392 In nearly aligned heterobilayers, electrons and holes can form bright interlayer excitons after the
393 charge transfer process^{28,40,83}. Rivera *et al.* observed 40% positive circular helicity from interlayer
394 exciton emission in WSe₂/MoSe₂ bilayer and measured a spin-valley lifetime of a few nanoseconds in
395 a time-resolved photoluminescence (TR-PL) study⁸⁴, as shown in Fig. 4c. The significantly longer
396 spin-valley lifetime of interlayer excitons can be understood by noting that the electron and hole
397 wavefunctions have much smaller overlap in interlayer excitons compared to intralayer excitons; they
398 will therefore have a weaker exchange interaction and exhibit slower recombination processes (both
399 radiative and non-radiative).

400

401 Recently, there has been increasing research interest in the nature of the interlayer exciton state and
402 the origin of the circular helicity of emission in nearly aligned heterostructures. Hsu *et al.* reported
403 negative circular helicity of interlayer exciton emission⁸⁵ in WSe₂/MoSe₂ bilayer, while Ciarrocchi *et al.*
404 and Hanbicki *et al.* observed two separate interlayer exciton emission peaks with opposite signs of
405 helicity^{86,87}. Meanwhile, various configurations of interlayer excitons have been proposed as the
406 emitting state, including spin singlet zero-momentum excitons^{84,88}, spin-triplet zero-momentum
407 excitons⁸⁹, and finite-momentum excitons^{90,91}. The rich set of observations originates in part from the
408 complex conduction band structure in the WSe₂/MoSe₂ bilayer, where electron states of spin up and
409 spin down, and at K and Q valleys, are all close in energy. Furthermore, the real-space distribution of
410 interlayer excitons when a moiré pattern is present can further modify the optical selection rules^{92,93}.
411 The exact mechanisms behind these interesting observations are yet to be fully understood.

412

413 An alternative approach involves using single-particle states, such as holes in WSe₂ to carry valley
414 information in the heterostructure. Because the momentum match between electrons and holes
415 (required for efficient exciton emission) is not relevant in this approach, a large-twist-angle bilayer is

416 preferred to separate electrons and holes in momentum space and further reduce their exchange
417 interaction and recombination rate. Kim *et al.* measured the spin-valley lifetime of holes in
418 WSe₂/MoS₂ heterostructures using circularly polarized pump-probe spectroscopy⁹⁴. Figure 4d shows
419 the measured decay dynamics of the total hole population and valley-polarized hole population in the
420 WSe₂ layer at a temperature of 10 K. Both the population lifetime and the spin-valley lifetime of holes
421 are around one microsecond, indicating that the decay of the spin-valley imbalance occurs primarily
422 through population loss. On the other hand, the valley polarization remains almost a constant for a
423 few microseconds, from which one can extract a valley depolarization time (or intervalley scattering
424 time) exceeding 40 μs. The other critical figure of merit, the conversion efficiency, was also
425 determined experimentally to be close to unity for valley-polarized holes⁹⁴. The nearly ideal
426 conversion efficiency is consistent with an interlayer charge transfer process that is far faster than
427 intervalley scattering processes.

428
429 The spin-valley lifetime of resident holes in the heterostructure can be further improved by tuning the
430 carrier concentration. Figure 4e summarizes the doping-dependent spin-valley lifetime of holes in a
431 WSe₂/WS₂ heterostructure (red circles), as compared to the population lifetime of holes (blue
432 triangles)⁹⁵. In charge neutral and electron-doped heterostructures, the spin-valley lifetime is similar to
433 the population lifetime; however, for hole-doping, the spin-valley lifetime becomes orders of
434 magnitude longer than the population lifetime. This doping dependence is a consequence of the
435 interlayer electron-hole recombination process, as shown in Fig. 4f,g. For electron-doped or charge
436 neutral heterostructures (Fig. 4f), all of the holes in WSe₂ are pump-generated excess holes.
437 Therefore, when hole population decays to zero due to interlayer electron-hole recombination, no
438 holes -- and certainly no valley-polarized holes -- remain in the WSe₂. The valley lifetime is then
439 limited by the lifetime of the total hole excess. On the other hand, if the original hole density is much
440 greater than the photo-generated density, excess electrons in WS₂ will recombine with holes from
441 both valleys of WSe₂ with nearly equal probability (Fig. 4g). Consequently, a pure spin-valley
442 imbalance (*i.e.*, equal excess and deficiency of holes in the K and K' valley) is generated, the lifetime
443 of which can be much longer than the population lifetime and has been found to exceed 20 μs.
444 Furthermore, the decrease of spin-valley imbalance is negligible during the population decay, and the
445 overall efficiency of this two-step conversion process can approach 100%⁹⁵.

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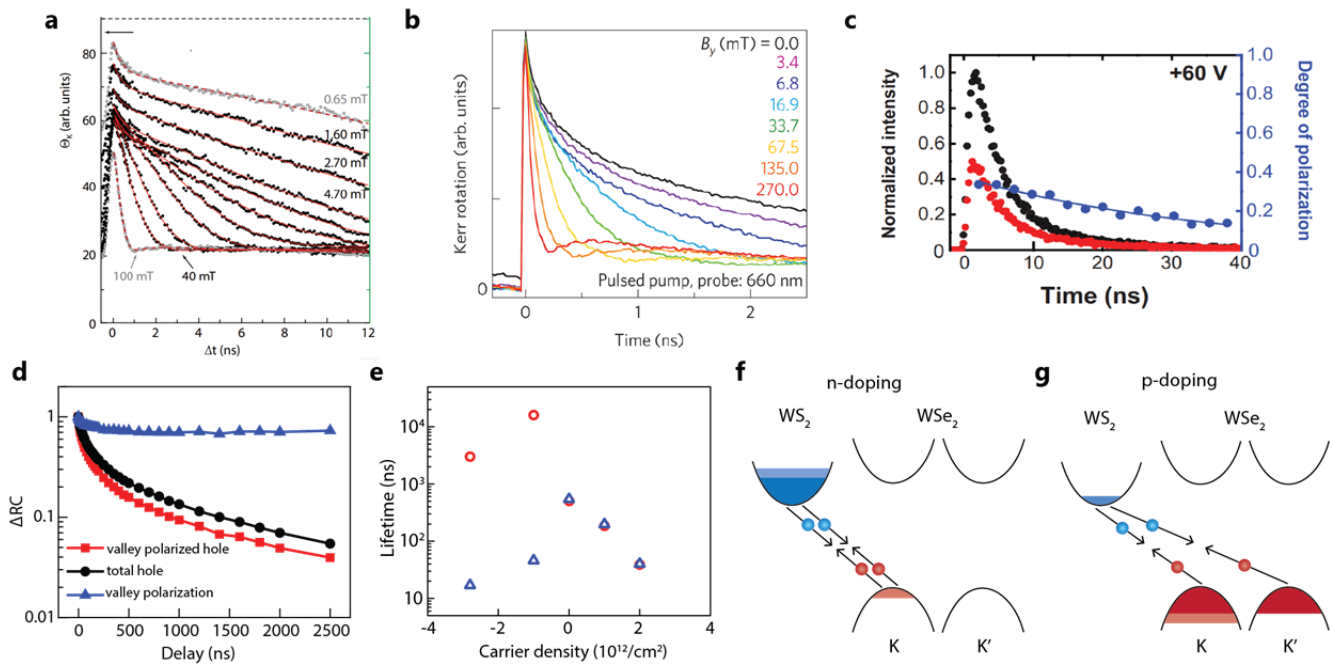


Fig. 4 Dynamics of spin-valley information carriers in TMDC

materials. (a-b) Spin-valley dynamics of trions (a)⁷² and resident electrons (b)⁸¹ in monolayer TMDC materials under different external magnetic fields as probed by time-resolved Kerr rotation measurements.

A spin-valley lifetime of few to tens of nanoseconds is observed at low fields. (c) Time-resolved photoluminescence in a nearly aligned WSe₂/MoSe₂ heterobilayer for the same (black) and opposite (red) circularly polarized excitation, revealing a valley lifetime of few nanoseconds⁸⁴.

(d) Decay dynamics of the valley-polarized hole population, the total hole population, and the valley polarization in a large-twist-angle WSe₂/MoS₂ heterostructure at charge neutrality⁹⁴.

(e) Summary of the hole population (blue) and spin-valley (red) lifetimes as a function of carrier concentration in a large-twist-angle WSe₂/WS₂ heterostructure⁹⁵.

(f-g) Schematic illustration of the interlayer electron-hole recombination process in electron-doped (f) and hole-doped (g) heterostructures.

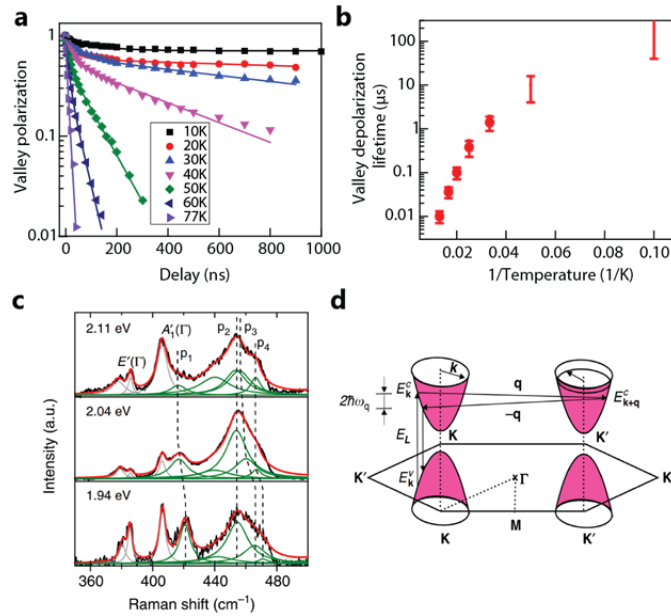
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469 Potential mechanism behind intervalley scattering

470 As discussed in the previous sections, the spin-valley lifetime can be limited either by the population
471 lifetime or by the intervalley scattering process. The first limitation can be removed for valley-polarized
472 holes in hole-doped heterostructures, making them promising candidates for spin-valley information
473 carriers. Furthermore, directly probing hole dynamics provides relatively clean information about the
474 decay mechanism owing to the simplicity of the valence band maximum. This configuration thus
475 provides a valuable platform for understanding intervalley scattering processes in TMDC materials.

476
477 Figure 5a shows the decay dynamics of valley polarization at different temperature for holes in
478 WSe₂/MoS₂ heterostructures, with the valley depolarization lifetime summarized in Fig. 5b (Ref. ⁹⁴).

479 The depolarization lifetime changes from 10 ns at 77 K to above 40 μ s at 10 K, which roughly follows
 480 a thermally activated rate: $\tau \sim \exp\left(\frac{\Delta}{k_B T}\right)$, with k_B denoting the Boltzmann constant and $\Delta \sim 20$ meV.
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 482



483
 484 **Fig. 5 Potential origin of intervalley scattering in WSe₂/MoS₂**
 485 heterostructures. (a) Temperature-dependent decay dynamics of valley
 486 polarization. (b) Temperature dependence of the intervalley scattering
 487 time. The process can be described as thermally activated⁹⁴. (c)
 488 Resonant Raman spectra of single-layer MoS₂ show particularly strong
 489 second-order Raman signals (peaks p1 to p4) for excitation around 2.0
 490 eV through a doubly resonant Raman (DRR) process involving K point
 491 phonons, as illustrated in (d)⁹⁶.
 492

493 The inter-valley scattering of holes in WSe₂ requires a large momentum change and a simultaneous
 494 spin flip. However, due to the strong spin-orbit coupling in WSe₂, the picture of electrons and holes
 495 with perfectly defined up and down spin states in the K and K' valleys, respectively, is not strictly valid
 496 for states away from the K and K' points. Therefore, inter-valley scattering of carriers near, but not
 497 exactly at K and K' points is allowed. This process, often designated as the Elliott-Yafet mechanism,
 498 has a characteristic temperature dependence of

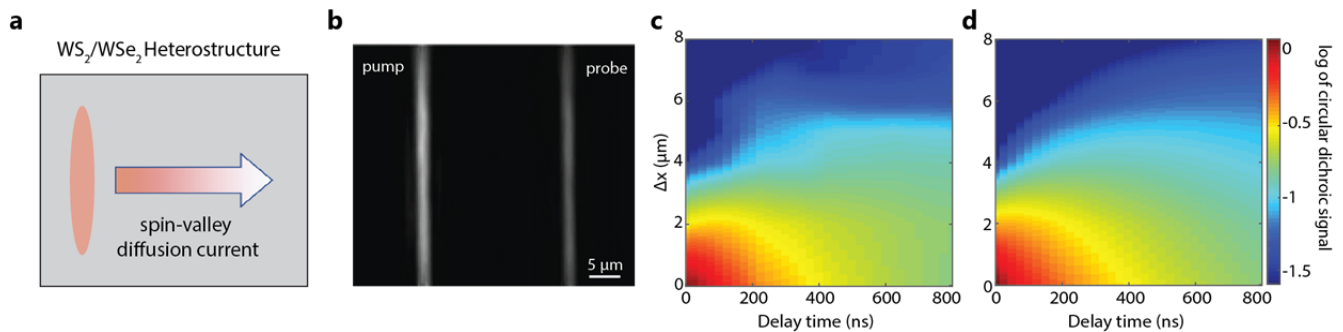
$$\tau_{EY}^{-1} \sim T^2 \tau_p^{-1},$$

499 where τ_p is the momentum scattering lifetime for a spin-conserving process. The T^2 -dependence
 500 originates from the fact that at higher temperature, thermally excited carriers are further away from
 501 band minima (K or K') and will therefore scatter more efficiently. However, the predicted T^2
 502 dependence does not describe the strong observed variation with temperature, which presumably
 503 reflects from the temperature dependence of τ_p^{-1} . Indeed, a thermally activated temperature
 504 dependence is expected for phonon-assisted intervalley scattering at low temperatures, and the
 505 experimental activation energy of ~ 20 meV agrees with WSe₂ phonon energy (at the K-point) required
 506 for intervalley scattering⁹⁶. Phonon-assisted intervalley scattering, accompanied by spin flip through
 507 the Elliott-Yafet mechanism, can thus account for the observed spin-valley depolarization of holes.
 508

509 The important role of K-point phonons in intervalley scattering is also supported by a recent resonant
 510 Raman study⁹⁶, which reveals second-order Raman signals (peaks p1 to p4 in Fig. 5c) assigned to K-
 511 point phonons through a doubly resonant Raman (DRR) scattering process. This observation
 512 suggests a strong interaction between charge carriers and K-point phonons, which dramatically
 513 enhances their second-order Raman signals through the DRR process illustrated in Fig. 5d.
 514

515 Spin-valley transport in TMDC heterostructures

516 The efficient generation of pure spin-valley imbalance in hole-doped WSe₂/WS₂ heterostructures
 517 provides a convenient way to create pure spin-valley current (Fig. 6a), which lies in the heart of spin-
 518 valley-tronic devices. Jin *et al.* performed space-and-time-resolved pump-probe spectroscopy to track
 519 the evolution of the spin-valley imbalance and to image the flow of pure spin-valley diffusion currents
 520 (Fig. 6b). Figure 6c shows experimental results for a hole-doped WS₂/WSe₂ heterostructure at an
 521 initial electrostatic hole doping of $p_0 = 1 \times 10^{12} / \text{cm}^2$. At zero time delay, the spin-valley imbalance
 522 matches the pump beam spatial profile (half-width of $\sim 1.5 \mu\text{m}$); the signal is negligible for a pump-
 523 probe separations greater than $3 \mu\text{m}$, as expected based on the convolution with probe spatial profile.
 524 At finite delay time, the spin-valley imbalance diffuses out of the excitation region, generating a pure
 525 spin-valley current. This leads to a strong decrease and increase of signal, respectively, in regions
 526 near and far away from the pump beam. The spin-valley current propagates to distances over $8 \mu\text{m}$
 527 within 800 ns. Such direct imaging of the experimental spin-valley current flow (Fig. 6c) allows us to
 528 determine important physical parameters by comparison with a diffusion-decay model (Fig. 6d). We
 529 infer a diffusion constant of $D = 0.2 \text{ cm}^2/\text{s}$, a spin-valley lifetime of $\tau = 20 \mu\text{s}$, and deduce a spin-valley
 530 diffusion length of $l = \sqrt{D\tau} = 20 \mu\text{m}$.
 531



532
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 534 Fig. 6 **Spin-valley transport in a vdW heterostructure**⁹⁵. (a) Optical
 535 excitation of pure spin-valley imbalance at the left edge of a device will
 536 create a pure spin-valley diffusion current flowing to the right without any
 537 associated charge current. (b) Experimental configuration for direct
 538 imaging of the spin-valley current flow with space-and-time resolved
 539 pump-probe spectroscopy using pump and probe beams focused to lines
 540 on the sample at defined spatial separation. (c-d) Experimentally
 541 measured spatio-temporal evolution of the pure valley imbalance in the
 542 heterostructure (c) and a simulation of the results using a diffusion-decay
 543 model (d) for an initial hole doping of $10^{12}/\text{cm}^2$.
 544

545 The efficient generation of spin-valley current with remarkably large current densities reflects the
 546 nearly ideal conversion of photogenerated excitons into a spin-valley imbalance. Still higher spin-

547 valley currents may be achievable in TMDC heterostructures by increasing the initial hole doping
548 level to enable stronger optical pumping, as well as by improving the device quality to enhance the
549 diffusivity. The long spin-valley lifetimes and diffusion lengths of valley-polarized holes in TMDC
550 heterostructures hold promise for the generation, transport, and detection of spin-valley information
551 and open exciting opportunities for the realization of future spintronic and valleytronic devices.
552

553 **Concluding remarks**

554 Despite the rapid progress in the study of excited-states in van der Waals heterostructures
555 summarized above, many outstanding questions remain in understanding charge transfer processes
556 and the spin and valley relaxation dynamics.
557

558 A complete picture of the underlying mechanism for the CT process in TMDC heterostructures
559 remains elusive. First, the mechanisms developed to date do not account for the Coulombic
560 interactions between electrons and holes, despite the existence of strongly bound excitons both in the
561 TMDC monolayers^{8,97,98} and in the heterostructures⁴⁰. Though one can argue that the excitonic states
562 are superpositions of the quasiparticle band states and, hence, are affected similarly, a quantitative
563 picture in which excitonic correlations are taken into account remains a theoretical challenge. In this
564 context, it would also be interesting to learn whether the dielectric environment of the heterostructure,
565 which affects the excitonic interactions, also significantly influences the rate and efficiency of CT
566 processes. In addition, the fact that changes in the dielectric screening of Coulombic interactions in
567 TMDCs modify the quasiparticle band structure may provide a route to test the role of the Q and Γ
568 valleys in CT processes. Second, the time-domain probes of CT to date have been limited in their by
569 the instrumental response function and have not generally yielded precise CT times. In addition,
570 optical measurements, with their limited spatial resolution, average over moiré patterns formed
571 between the two layers. This may lead to a washing out of predicted trends for CT times. Overcoming
572 these limitations by improved temporal resolution and/or spatial resolution (such as through near-field
573 techniques) would provide important experimental information to inform and test further theoretical
574 models.
575

576 Similarly many outstanding questions exist regarding spin and valley dynamics in TMDC
577 heterostructures. Much more work is required to understand fully the intrinsic spin and valley
578 dynamics in TMDC heterostructures and the dependence of the dynamics on the constituent TMDC
579 materials, their relative crystallographic alignment, and their stacking order in multilayers. It remains,
580 for example, unclear what factors define the ultimate limit for the spin-valley lifetime in TMDC
581 heterostructures; also unknown is the role played by defects, edges, and grain boundaries, as well as
582 possible effects from large-period moiré superlattices formed in TMDC heterostructures with small
583 twist angles.
584

586 **References**

587

- 588 1 Hunt, B. *et al.* Massive Dirac Fermions and Hofstadter Butterfly in a van der Waals
589 Heterostructure. *Science* **340**, 1427-1430, (2013).
- 590 2 Ponomarenko, L. A. *et al.* Cloning of Dirac fermions in graphene superlattices. *Nature* **497**,
591 594-597, (2013).
- 592 3 Dean, C. R. *et al.* Hofstadter's butterfly and the fractal quantum Hall effect in moire
593 superlattices. *Nature* **497**, 598-602, (2013).
- 594 4 Spanton, E. M. *et al.* Observation of fractional Chern insulators in a van der Waals
595 heterostructure. *Science* **360**, 62-66, (2018).
- 596 5 Cao, Y. *et al.* Unconventional superconductivity in magic-angle graphene superlattices. *Nature*
597 **556**, 43-+, (2018).
- 598 6 Cao, Y. *et al.* Correlated insulator behaviour at half-filling in magic-angle graphene
599 superlattices. *Nature* **556**, 80-+, (2018).
- 600 7 Chen, G. *et al.* Gate-Tunable Mott Insulator in Trilayer Graphene-Boron Nitride Moiré
601 Superlattice. *Arxiv*, 1803.01985, (2018).
- 602 8 Chernikov, A. *et al.* Exciton Binding Energy and Nonhydrogenic Rydberg Series in Monolayer
603 WS₂. *Phys Rev Lett* **113**, (2014).
- 604 9 Ye, Z. L. *et al.* Probing excitonic dark states in single-layer tungsten disulphide. *Nature* **513**,
605 214-218, (2014).
- 606 10 Xiao, D., Liu, G. B., Feng, W. X., Xu, X. D. & Yao, W. Coupled Spin and Valley Physics in
607 Monolayers of MoS₂ and Other Group-VI Dichalcogenides. *Phys Rev Lett* **108**, (2012).
- 608 11 Xu, X. D., Yao, W., Xiao, D. & Heinz, T. F. Spin and pseudospins in layered transition metal
609 dichalcogenides. *Nat Phys* **10**, 343-350, (2014).
- 610 12 Komsa, H. P. & Krasheninnikov, A. V. Electronic structures and optical properties of realistic
611 transition metal dichalcogenide heterostructures from first principles. *Phys Rev B* **88**, (2013).
- 612 13 Gong, C. *et al.* Band alignment of two-dimensional transition metal dichalcogenides:
613 Application in tunnel field effect transistors. *Appl Phys Lett* **103**, (2013).
- 614 14 Terrones, H., Lopez-Urias, F. & Terrones, M. Novel hetero-layered materials with tunable
615 direct band gaps by sandwiching different metal disulfides and diselenides. *Sci Rep-Uk* **3**,
616 (2013).
- 617 15 Stormer, H. L., Dingle, R., Gossard, A. C., Wiegmann, W. & Sturge, M. D. 2-Dimensional
618 Electron-Gas at a Semiconductor-Semiconductor Interface. *Solid State Commun* **29**, 705-709,
619 (1979).
- 620 16 Bellus, M. Z. *et al.* Type-I van der Waals heterostructure formed by MoS₂ and ReS₂
621 monolayers. *Nanoscale Horiz* **2**, 31-36, (2017).
- 622 17 Chiu, M. H. *et al.* Determination of band alignment in the single-layer MoS₂/WSe₂
623 heterojunction. *Nat Commun* **6**, (2015).
- 624 18 Ozcelik, V. O., Azadani, J. G., Yang, C., Koester, S. J. & Low, T. Band alignment of two-
625 dimensional semiconductors for designing heterostructures with momentum space matching.
626 *Phys Rev B* **94**, (2016).
- 627 19 Hong, X. P. *et al.* Ultrafast charge transfer in atomically thin MoS₂/WS₂ heterostructures. *Nat*
628 *Nanotechnol* **9**, 682-686, (2014).
- 629 20 Wang, K. *et al.* Interlayer Coupling in Twisted WSe₂/WS₂ Bilayer Heterostructures Revealed
630 by Optical Spectroscopy. *Acs Nano* **10**, 6612-6622, (2016).
- 631 21 Kozawa, D. *et al.* Evidence for Fast Interlayer Energy Transfer in MoSe₂/WS₂
632 Heterostructures. *Nano Lett* **16**, 4087-4093, (2016).
- 633 22 Zhang, C. X. *et al.* Systematic study of electronic structure and band alignment of monolayer
634 transition metal dichalcogenides in Van der Waals heterostructures. *2d Mater* **4**, (2017).

- 635 23 Debbichi, L., Eriksson, O. & Lebegue, S. Electronic structure of two-dimensional transition
636 metal dichalcogenide bilayers from ab initio theory. *Phys Rev B* **89**, (2014).
- 637 24 Zhang, J. F., Xie, W. Y., Zhao, J. J. & Zhang, S. B. Band alignment of two-dimensional lateral
638 heterostructures. *2d Mater* **4**, (2017).
- 639 25 Qiu, D. Y., da Jornada, F. H. & Louie, S. G. Screening and many-body effects in two-
640 dimensional crystals: Monolayer MoS₂. *Phys Rev B* **93**, (2016).
- 641 26 Wilson, N. R. *et al.* Determination of band offsets, hybridization, and exciton binding in 2D
642 semiconductor heterostructures. *Sci Adv* **3**, (2017).
- 643 27 Hill, H. M., Rigosi, A. F., Rim, K. T., Flynn, G. W. & Heinz, T. F. Band Alignment in MoS₂/WS₂
644 Transition Metal Dichalcogenide Heterostructures Probed by Scanning Tunneling Microscopy
645 and Spectroscopy. *Nano Lett* **16**, 4831-4837, (2016).
- 646 28 Ceballos, F., Bellus, M. Z., Chiu, H. Y. & Zhao, H. Ultrafast Charge Separation and Indirect
647 Exciton Formation in a MoS₂-MoSe₂ van der Waals Heterostructure. *Acs Nano* **8**, 12717-
648 12724, (2014).
- 649 29 Heo, H. *et al.* Interlayer orientation-dependent light absorption and emission in monolayer
650 semiconductor stacks. *Nat Commun* **6**, (2015).
- 651 30 Zhu, H. M. *et al.* Interfacial Charge Transfer Circumventing Momentum Mismatch at Two-
652 Dimensional van der Waals Heterojunctions. *Nano Lett* **17**, 3591-3598, (2017).
- 653 31 Ji, Z. H. *et al.* Robust Stacking-Independent Ultrafast Charge Transfer in MoS₂/WS₂ Bilayers.
654 *Acs Nano* **11**, 12020-12026, (2017).
- 655 32 Chen, H. L. *et al.* Ultrafast formation of interlayer hot excitons in atomically thin MoS₂/WS₂
656 heterostructures. *Nat Commun* **7**, (2016).
- 657 33 Ceballos, F., Ju, M. G., Lane, S. D., Zeng, X. C. & Zhao, H. Highly Efficient and Anomalous
658 Charge Transfer in van der Waals Trilayer Semiconductors. *Nano Lett* **17**, 1623-1628, (2017).
- 659 34 Lee, C.-H. *et al.* Atomically thin p-n junctions with van der Waals heterointerfaces. *Nat*
660 *Nanotechnol* **9**, 676, (2014).
- 661 35 Rigos, A. F., Hill, H. M., Li, Y. L., Chernikov, A. & Heinz, T. F. Probing Interlayer Interactions in
662 Transition Metal Dichalcogenide Heterostructures by Optical Spectroscopy: MoS₂/WS₂ and
663 MoSe₂/WSe₂. *Nano Lett* **15**, 5033-5038, (2015).
- 664 36 Pan, S. D., Ceballos, F., Bellus, M. Z., Zereshki, P. & Zhao, H. Ultrafast charge transfer
665 between MoTe₂ and MoS₂ monolayers. *2d Mater* **4**, (2017).
- 666 37 Xu, W. G. *et al.* Correlated fluorescence blinking in two-dimensional semiconductor
667 heterostructures. *Nature* **541**, 62+, (2017).
- 668 38 Ceballos, F., Bellus, M. Z., Chiu, H. Y. & Zhao, H. Probing charge transfer excitons in a
669 MoSe₂-WS₂ van der Waals heterostructure. *Nanoscale* **7**, 17523-17528, (2015).
- 670 39 Li, Y. Y. *et al.* Ultrafast Interlayer Electron Transfer in Incommensurate Transition Metal
671 Dichalcogenide Homobilayers. *Nano Lett* **17**, 6661-6666, (2017).
- 672 40 Rivera, P. *et al.* Observation of long-lived interlayer excitons in monolayer MoSe₂-WSe₂
673 heterostructures. *Nat Commun* **6**, (2015).
- 674 41 Robert, C. *et al.* Exciton radiative lifetime in transition metal dichalcogenide monolayers. *Phys*
675 *Rev B* **93**, (2016).
- 676 42 Xu, W. S. *et al.* Determining the Optimized Interlayer Separation Distance in Vertical Stacked
677 2D WS₂:hBN:MoS₂ Heterostructures for Exciton Energy Transfer. *Small* **14**, (2018).
- 678 43 Mak, K. F., Lee, C., Hone, J., Shan, J. & Heinz, T. F. Atomically Thin MoS₂: A New Direct-Gap
679 Semiconductor. *Phys Rev Lett* **105**, (2010).
- 680 44 Splendiani, A. *et al.* Emerging Photoluminescence in Monolayer MoS₂. *Nano Lett* **10**, 1271-
681 1275, (2010).
- 682 45 Zhu, X. Y. *et al.* Charge Transfer Excitons at van der Waals Interfaces. *J Am Chem Soc* **137**,
683 8313-8320, (2015).
- 684 46 Wang, H. *et al.* The role of collective motion in the ultrafast charge transfer in van der Waals
685 heterostructures. *Nat Commun* **7**, (2016).
- 686 47 Zhang, J. *et al.* Interlayer-State-Coupling Dependent Ultrafast Charge Transfer in MoS₂/WS₂
687 Bilayers. *Adv Sci* **4**, (2017).

688 48 Long, R. & Prezhdo, O. V. Quantum Coherence Facilitates Efficient Charge Separation at a
689 MoS₂/MoSe₂ van der Waals Junction. *Nano Lett* **16**, 1996-2003, (2016).

690 49 Li, L. Q., Long, R. & Prezhdo, O. V. Charge Separation and Recombination in Two-
691 Dimensional MoS₂/WS₂: Time-Domain ab Initio Modeling. *Chem Mater* **29**, 2466-2473, (2017).

692 50 Kang, J., Li, J. B., Li, S. S., Xia, J. B. & Wang, L. W. Electronic Structural Moire Pattern Effects
693 on MoS₂/MoSe₂ 2D Heterostructures. *Nano Lett* **13**, 5485-5490, (2013).

694 51 Tong, Q. J. *et al.* Topological mosaics in moire superlattices of van der Waals heterobilayers.
695 *Nat Phys* **13**, 356-362, (2017).

696 52 Wang, Y., Wang, Z., Yao, W., Liu, G. B. & Yu, H. Y. Interlayer coupling in commensurate and
697 incommensurate bilayer structures of transition-metal dichalcogenides. *Phys Rev B* **95**, (2017).

698 53 Zheng, Q. J. *et al.* Phonon-Assisted Ultrafast Charge Transfer at van der Waals
699 Heterostructure Interface. *Nano Lett* **17**, 6435-6442, (2017).

700 54 Jin, C. H. *et al.* On Optical Dipole Moment and Radiative Recombination Lifetime of Excitons
701 in WSe₂. *Adv Funct Mater* **27**, (2017).

702 55 Zhang, X. X., You, Y. M., Zhao, S. Y. F. & Heinz, T. F. Experimental Evidence for Dark
703 Excitons in Monolayer WSe₂. *Phys Rev Lett* **115**, (2015).

704 56 Korn, T., Heydrich, S., Hirmer, M., Schmutzler, J. & Schuller, C. Low-temperature photocarrier
705 dynamics in monolayer MoS₂. *Appl Phys Lett* **99**, (2011).

706 57 Amani, M. *et al.* Near-unity photoluminescence quantum yield in MoS₂. *Science* **350**, 1065-
707 1068, (2015).

708 58 Moody, G. *et al.* Intrinsic homogeneous linewidth and broadening mechanisms of excitons in
709 monolayer transition metal dichalcogenides. *Nat Commun* **6**, (2015).

710 59 Lagarde, D. *et al.* Carrier and Polarization Dynamics in Monolayer MoS₂. *Phys Rev Lett* **112**,
711 (2014).

712 60 Wang, Q. S. *et al.* Valley Carrier Dynamics in Mono layer Molybdenum Disulfide from Helicity-
713 Resolved Ultrafast Pump-Probe Spectroscopy. *ACS Nano* **7**, 11087-11093, (2013).

714 61 Hao, K. *et al.* Direct measurement of exciton valley coherence in monolayer WSe₂. *Nat Phys*
715 **12**, 677+, (2016).

716 62 Mai, C. *et al.* Many-Body Effects in Valleytronics: Direct Measurement of Valley Lifetimes in
717 Single-Layer MoS₂. *Nano Lett* **14**, 202-206, (2014).

718 63 Zhu, C. R. *et al.* Exciton valley dynamics probed by Kerr rotation in WSe₂ monolayers. *Phys*
719 *Rev B* **90**, (2014).

720 64 Mak, K. F., He, K. L., Shan, J. & Heinz, T. F. Control of valley polarization in monolayer MoS₂
721 by optical helicity. *Nat Nanotechnol* **7**, 494-498, (2012).

722 65 Zeng, H. L., Dai, J. F., Yao, W., Xiao, D. & Cui, X. D. Valley polarization in MoS₂ monolayers
723 by optical pumping. *Nat Nanotechnol* **7**, 490-493, (2012).

724 66 Cao, T. *et al.* Valley-selective circular dichroism of monolayer molybdenum disulphide. *Nat*
725 *Commun* **3**, (2012).

726 67 Glazov, M. M. *et al.* Exciton fine structure and spin decoherence in monolayers of transition
727 metal dichalcogenides. *Phys Rev B* **89**, (2014).

728 68 Yu, T. & Wu, M. W. Valley depolarization due to intervalley and intravalley electron-hole
729 exchange interactions in monolayer MoS₂. *Phys Rev B* **89**, (2014).

730 69 Yu, H. Y., Liu, G. B., Gong, P., Xu, X. D. & Yao, W. Dirac cones and Dirac saddle points of
731 bright excitons in monolayer transition metal dichalcogenides. *Nat Commun* **5**, (2014).

732 70 Maialle, M. Z., Silva, E. A. D. E. & Sham, L. J. Exciton Spin Dynamics in Quantum-Wells. *Phys*
733 *Rev B* **47**, 15776-15788, (1993).

734 71 Hao, K. *et al.* Trion valley coherence in monolayer semiconductors. *2d Mater* **4**, (2017).

735 72 Volmer, F. *et al.* Intervalley dark trion states with spin lifetimes of 150 ns in WSe₂. *Phys Rev B*
736 **95**, (2017).

737 73 Plechinger, G. *et al.* Trion fine structure and coupled spin-valley dynamics in monolayer
738 tungsten disulfide. *Nat Commun* **7**, (2016).

739 74 Zhang, X. X. *et al.* Magnetic brightening and control of dark excitons in monolayer WSe₂. *Nat*
740 *Nanotechnol* **12**, 883+, (2017).

741 75 Jiang, C. Y. *et al.* Microsecond dark-exciton valley polarization memory in two-dimensional
742 heterostructures. *Nat Commun* **9**, (2018).

743 76 You, Y. M. *et al.* Observation of biexcitons in monolayer WSe₂. *Nat Phys* **11**, 477-U138,
744 (2015).

745 77 Chen, S., Goldstein, T., Taniguchi, T., Watanabe, K. & Yan, J. Coulomb-bound four- and five-
746 particle valleytronic states in an atomically-thin semiconductor. *Arxiv*, 1802.10247, (2018).

747 78 Dey, P. *et al.* Gate-Controlled Spin-Valley Locking of Resident Carriers in WSe₂ Monolayers.
748 *Phys Rev Lett* **119**, (2017).

749 79 Song, X. L., Xie, S. E., Kang, K., Park, J. & Sih, V. Long-Lived Hole Spin/Valley Polarization
750 Probed by Kerr Rotation in Monolayer WSe₂. *Nano Lett* **16**, 5010-5014, (2016).

751 80 Hsu, W. T. *et al.* Optically initialized robust valley-polarized holes in monolayer WSe₂. *Nat*
752 *Commun* **6**, (2015).

753 81 Yang, L. Y. *et al.* Long-lived nanosecond spin relaxation and spin coherence of electrons in
754 monolayer MoS₂ and WS₂. *Nat Phys* **11**, 830-U187, (2015).

755 82 Fang, H. *et al.* Strong interlayer coupling in van der Waals heterostructures built from single-
756 layer chalcogenides. *P Natl Acad Sci USA* **111**, 6198-6202, (2014).

757 83 Nagler, P. *et al.* Interlayer exciton dynamics in a dichalcogenide monolayer heterostructure. *2d*
758 *Mater* **4**, (2017).

759 84 Rivera, P. *et al.* Valley-polarized exciton dynamics in a 2D semiconductor heterostructure.
760 *Science* **351**, 688-691, (2016).

761 85 Hsu, W. T. *et al.* Negative circular polarization emissions from WSe₂/MoSe₂ commensurate
762 heterobilayers. *Nat Commun* **9**, (2018).

763 86 Hanbicki, A. T. *et al.* Double Indirect Interlayer Exciton in a MoSe₂/WSe₂ van der Waals
764 Heterostructure. *Arxiv*, 1802.05310, (2018).

765 87 Ciarrocchi, A. *et al.* Control of interlayer excitons in two-dimensional van der Waals
766 heterostructures. *Arxiv*, 1803.06405, (2018).

767 88 Yu, H. Y., Wang, Y., Tong, Q. J., Xu, X. D. & Yao, W. Anomalous Light Cones and Valley
768 Optical Selection Rules of Interlayer Excitons in Twisted Heterobilayers. *Phys Rev Lett* **115**,
769 (2015).

770 89 Yu, H. Y., Liu, B. L. & Yao, W. Brightened spin-triplet interlayer excitons and optical selection
771 rules in van der Waals heterobilayers. 1803.01292, (2018).

772 90 Miller, B. *et al.* Long-Lived Direct and Indirect Interlayer Excitons in van der Waals
773 Heterostructures. *Nano Lett* **17**, 5229-5237, (2017).

774 91 Kunstmann, J. *et al.* Momentum-space indirect interlayer excitons in transition-metal
775 dichalcogenide van der Waals heterostructures. *Nat Phys*, (2018).

776 92 Wu, F. C., Lovorn, T. & MacDonald, A. H. Theory of optical absorption by interlayer excitons in
777 transition metal dichalcogenide heterobilayers. *Phys Rev B* **97**, (2018).

778 93 Yu, H. Y., Liu, G. B., Tang, J. J., Xu, X. D. & Yao, W. Moire excitons: From programmable
779 quantum emitter arrays to spin-orbit-coupled artificial lattices. *Sci Adv* **3**, (2017).

780 i94 Kim, J. *et al.* Observation of ultralong valley lifetime in WSe₂/MoS₂ heterostructures. *Sci Adv*
781 **3**, (2017).

782 95 Jin, C. H. *et al.* Imaging of Pure Spin-Valley Diffusion Current in WS₂/WSe₂ Heterostructures.
783 *Science*, in press, (2018).

784 96 Carvalho, B. R. *et al.* Intervalley scattering by acoustic phonons in two-dimensional MoS₂
785 revealed by double-resonance Raman spectroscopy. *Nat Commun* **8**, (2017).

786 97 Hill, H. M. *et al.* Observation of Excitonic Rydberg States in Monolayer MoS₂ and WS₂ by
787 Photoluminescence Excitation Spectroscopy. *Nano Lett* **15**, 2992-2997, (2015).

788 98 He, K. L. *et al.* Tightly Bound Excitons in Monolayer WSe₂. *Phys Rev Lett* **113**, (2014).

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